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An Atomic-Resolution Study of Overlayer Formation and Interfacial Mixing in the Interaction of Phosphorus with Si(001)

by

Y. Wang, X. Chen, and R.J. Hamers

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An Atomic-Resolution Study of Overlayer Formation and Interfacial Mixing in the Interaction of Phosphorus with Si(001)

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Abstract

Scanning Tunneling Microscopy (STM), tunneling spectroscopy, and Auger electron spectroscopy have

heterodimers and the simultaneous disappearance of are used to study the equilibrium between Si=Si, Si-P, show that surface phosphorus atoms readily displace and P-P dimers, showing that the surface is a nearly decomposition of phosphine (PH3). The STM images overall surface morphology through the formation numbers of line defects which act as a strain relief the strain-induced line defects. Counting statistics terminated with P-P dimers, but also contain large mechanism. STM images are used to determine the lower phosphorus coverage, the STM experiments revealing the formation of large numbers of Si-P anisotropic islands and very rough step edges. At Si from the substrate, dramatically changing the symmetry of these line defects and are compared with model defect structures. Line defects in the dimension, such that the phosphorus-terminated been used to study the formation of phosphorusterminated silicon(001) surfaces by the thermal are able to identify Si and P atoms individually, phosphorus-terminated surface both nucleate islands and also constrain their growth in one surfaces contain large numbers of extremely of large numbers of islands and an extreme roughening of step edges. The surfaces are random alloy with a slight non-statistical

are used to study the equilibrium between Si=Si, Si-P, and P-P dimers, showing that the surface is a nearly random alloy with a slight non-statistical preference for formation of the Si-P heterodimer. The geometric and electronic properties of Si=Si, Si-P, and P-P dimers on Si(001) are discussed on the basis of tunneling microscopy and tunneling spectroscopy measurements.

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Introduction

Phosphorus is a common dopant used in the fabrication of n-type silicon, usually introduced in chemical vapor deposition (CVD) processes through the decomposition of phosphine, PH3. Several studies have shown that during the growth of silicon from chemical precursors such as silane, small amounts of PH3 strongly influence the growth kinetics, decreasing the Si growth rate far out of proportion to the PH3 concentration in the gas stream.

In the last several years there has been a great interest in the use of surfactants to control surface stress in the growth of heterointerfaces, most notably the growth of Ge on Si using As or Sb as surfactants. While deposition of Ge on pure Si produces three-dimensional island growth after three monolayers due to the 4.3% lattice mismatch between Ge and Si, Copel and co-workers⁶, ⁷ have shown that incorporation of arsenic or antimony as surfactants during growth stabilizes the surface against three-dimensional growth by controlling the surface tension. Using this surfactant growth technique, they were able to grow Ge overlayers of

arbitrary thickness on Si without forming islands. Since phosphorus is a group V element isoelectronic with both As and Sb, phosphorus might be expected to show similar surfactant properties.

adsorption depends on how the surface free energy varies as a function of coverage. In the analysis of group V elements, it should be possible with STM to dimers can form atop a Si substrate or can displace recently suggested that heterodimers may play an both in the use of adlayers of group V elements as elements into the crystal lattice as dopants. In the silicon to form As-As dimers in the surface plane, Interfacial mixing plays an important role widely considered. However, Yu and Oshiyama . heterodimers such as Si-P or Si-As have not been important role in surface doping. Because of the particular case of arsenic, it is known that As-As tendency toward displacive adsorption vs. simple Tromp, as well as most other studies of surfactant surfactants as well as the incorporation of these depending on the preparation conditions. 13, 14 Tromp and co-workers have suggested that the different electronic structures of group IV and growth, it is usually assumed that the surface contains As-As dimers and Si-Si dimers, but

investigate the composition of such overlayers and to determine whether in fact such heterodimers are present and, if present, whether they play any role in the surface energetics or growth kinetics.

than 675 Kelvin (the desorption temperature of H2), the surface region until the surface P coverage was diffraction (LEED) to investigate the adsorption and exposure to PH3 produced a phosphorus-terminated monolayer, and which showed a streaky (2x1) LEED found that phosphorus was not effective in doping performed the most extensive studies, using Auger surface. They found that at temperatures higher thereby decreasing CVI) growth rates. They also Despite its widespread use as a dopant and phosphorus on Si(001) has not been extensively passivated it against further chemical reaction, phosphorus dimers formed on the surface and studied. Yu, Meyerson, and co-workers^{5, 16, 17} dissociation of phosphine (PH3) on the Si(001) spectroscopy(XPS), temperature-programmed surface with a coverage of approximately one electron spectroscopy, X-ray photoelectron desorption (TPD), and low-energy electron potential as a surfactant, the adsorption of pattern. They proposed that phosphorus-

very close to one monolayer, at which point the Fermi level suddenly jumped by 0.45 eV to near the conduction band. At sufficiently high temperatures they found that phosphorus desorbed from the surface, mostly in the form of P2.

In this work, we report on STM investigations of the atomic structure of phosphorus overlayers on silicon formed by the thermal decomposition of PH3. structures of Si=Si, Si-P, and P-P dimers, we are able basis. For the P-Si system, we find that the chemical alloys of P and Si, but with a slightly non-statistical composition of surfaces produced by adsorption of produced by desorption of phosphorus from the Ppreference for formation of the P-Si heterodimer. terminated surface both consist of nearly random phosphorus onto clean Si(001) as well as surfaces We find that phosphorus readily displaces silicon from the substrate, resulting in extensive island composition of the surface on an atom-by-atom coverages the dominant chemical specie on the surface is the Si-P heterodimer. Our results also formation. Because of the differing electronic to identify these three kinds of dimers on the As a result, over a wide range of phosphorus surface and are able to study the chemical

show the stress-induced line defects occur in this system at saturation coverage where P-P dimers dominate, but not at lower coverage where Si-P dimers dominate. Based on the symmetry of the line defects with respect to the underlying crystal lattice, we show that these line defects cannot be the same defects recently proposed by Tersoff 12 to account for similar-appearing line defects in the growth of strained Ge films on Si(001) We also show that the line defects present on the P-terminated surface significantly modify the mechanisms of island nucleation and growth.

Experimental

ultrahigh vacuum chamber with a base pressure of below 1x10⁻¹⁰ torr. Wafers of n-type Si(001) oriented to within 0.5° obtained from Wacker Chemitronic were used. Samples were prepared by first degassing for about 10 hours at 900 Kelvin, then annealing to 1400 Kelvin for several seconds on a separate heating stage. Temperatures were measured using an infrared pyrometer. After allowing 30 min for cooling to room temperature, samples were transferred to the STM and imaged to

ensure that the starting surfaces were clean and well ordered. Average terrace widths of nearly 1,000 Å were measured in the STM, with atomic-resolution images showing surface structure commensurate with previous studies.

controlled temperature of approximately 820 Kelvin, operation using tunneling currents (It) of less than then cooled, moved back to the STM tunneling stage, (probing occupied surface states) and +3V (probing measurements were performed by interrupting the Phosphorus overlayers were prepared from the in-1.0 nA with the sample bias (Vsample) between -3V silicon surface to ensure that the starting surfaces and imaged at room temperature. All images were PH3(99.999%, Matheson). After imaging the clean and then exposed to PH3 admitted into the vacuum chamber through a leak valve. The samples were STM feedback system and ramping the sample-tip were clean and well-ordered, the silicon samples unoccupied surface states), as indicated in the bias while keeping the sample-tip separation were moved to the heating stage, heated to a acquired in the "constant-current" mode of situ thermal decomposition of high purity figure captions. Tunneling spectroscopy constant

Results

a) Structure and properties of phosphorus overlayers near monolayer coverage

 $1 L = 10^{-6}$ torr-seconds), and then cooling the sample hydrogen from the PH3 decomposition. Between 675 successively higher resolution of surfaces prepared 825 Kelvin, exposing the surface to 10-7 torr PH3 for image includes parts of five different terraces, with 200 seconds (total exposure of 20 Langmuirs, where decomposition on Si(001) is temperature dependent. conditions. Figure la shows an STM image of a 1200 by heating a clean, well-ordered Si(001) surface to approximately 0.25, due to the presence of surface overall surface morphology on a large scale. This Yu and co-workers showed that phosphine maximum of one monolayer near 825 Kelvin, and upper right and the lowest terrace, labeled "5", at K and 825 K the coverage increases, reaching a Å x 1150 Å region of the surface, revealing the the highest terrace, labeled "1" at the extreme decreasing again at T>875 K due to phosphorus At low temperatures the maximum coverage is desorption. Figures 1-3 show STM images at back to 300 Kelvin under ultrahigh vacuum

islands on the terraces, whereas the starting surface the 90 degree rotation of the crystal structure across consisted of flat terraces with flat terraces separated extensions and deep invaginations extending nearly slightly asymmetric STM tip makes the defects more morphological change produced by PH3 exposure is morphological change produced by PH3 exposure is apparent on some terraces and not on others due to exposure to PH3 induces three dramatic changes in apparent in fig. 1a on terraces 1 and 3, these line visible on terraces 1 and 3. These line defects will which appear as dark, narrow lines, particularly the creation of very long, 1-dimensional defects lower left. Examination of this image shows that by monatomic steps. The second morphological the creation of a large number of long, narrow change created by PH3 exposure is the extreme the width of the individual terraces. The third roughening of the step edges, with fingerlike be denoted as "LD", and several LD defects are defects can be observed on all terraces, but a indicated in fig. 1a. Although they are most the overall surface morphology. The first a monatomic step on Si(001).

some cases the islands are only the width of a single also asymmetric in the same direction as the islands strings" ("DS") and have indicated several examples extend only one atomic layer deep. These holes are (easily visible at upper right and at lower left) are difficult to identify a single "substrate" terrace. In intersection of two islands under conditions where dimer strings always have a line defect on one side atoms do not diffuse over the islands to "fill in" the hole. One example of such as hole is labeled "H" in several hundred Angstroms, the terrace structure becomes dominated by the islands, and it becomes always bounded on both sides by line defects. The along their long dimension, while larger islands irrespective of the size of the island. Individual islands, particularly the islands visible at upper in fig. 1b. In this figure it can be seen that the apparent holes in the surface, which typically and are most likely produced by the imperfect right, are bounded on their long sides by line large number of very asymmetric islands also dimer row. We refer to such islands as "dimer results in the formation of a large number of Fig. 1b shows that on distance scales of defects in the underlying layer. This is true

involve substitution of phosphorus for silicon in the edge. This indicates that the observed structures are outermost atomic layer. 13, 14 While in principle the by obtaining STM images on extremely flat surfaces leads to the spontaneous formation of small regions manner similar to that of the clean Si(001) surface. fig. 1b. Finally, we note that the extreme degree of and measuring the fractional island coverage, this such as fig. 1-2, we find that comparatively smooth amount of phosphorus ejection could be calculated separations of nearly 1,000 Å, the extremely rough of double-height steps, several examples of which is not possible because even on our extremely flat step roughing and island formation in some cases step edges alternate with rougher step edges, in a rows run perpendicular to the double-height step edges, and at double steps (as in fig. 1b) the dimer not simply a phosphorus adlayer, but in all cases are labeled "2S" in fig. 1b. In large-scale images exposed surfaces run parallel to the smooth step In all cases, we find that the dimer rows of PH3step edges makes it impossible to distinguish starting surfaces, which have average step "islands" from step-edge roughness.

at resolution of fig. 2a and 2b, we are able to observe in previously been observed in silicon homocpitaxy. defects can be observed on both the upper and lower occasional shifts in direction. An examination of the troughs between dimer rows on opposite sides of the terraces. On the larger, lower terrace these defects example of line defects in which the dimer rows on defects are not the "antiphase" defects which have in almost all instances the dimer rows on each side labeled 1-3, with the highest terrace (1) consisting line defect are collinear). This shows that the line dimer rows on each side of these line defects (most surfaces prepared in the same manner as in fig. 1, easily visible by looking at the islands) shows that successively higher resolution. Fig. 2a consists of structure of the surface unit cell. In fig. 2a, rows two terraces. In fig. 2b, three planes are visible, of the line defect have the same phase (i.e., the can be observed on the upper terrace, and line extend for hundreds of Angstroms with only In fig. 2b at upper left, we can see a good more detail the structure of the line defects, Figures 2a and 2b show images of the nucleation site for islands, and the internal of only two rows of dimerized atoms. At the

opposite sides are in phase ("IPLD") and out-of-phase ("OPLD"); these regions are necessarily separated by an "AP1" antiphase boundary which involves a translational shift of 3.84 Å parallel to the dimer bond direction (perpendicular to the dimer rows), and appears similar to the AP1 boundaries observed in silicon homoepitxy.

22, 23 The in-phase and out-of-phase line defects appear to have similar structures, and we will henceforth simply refer to them collectively as line defects ("LD").

As noted earlier, these line defects appear to act as nucleation sites for next-year growth. In fig. 2a, there are several examples in which dimer strings have nucleated adjacent to (but not on top of) line defects. The dimer string at bottom right in fig. 2a is bounded on both sides by line defects. Similarly, at higher resolution fig. 2b shows that the two dimer rows constituting the topmost terrace are bounded on both sides by line defects. Surprisingly, in fig. 2a we see that the dimer strings at bottom right and at left center appear to crossed the gap between different islands, forming a bridge only a single dimer wide between the islands.

two atoms constituting a single dimer have different probability for each side), and appear as though the shared between the two Si atoms. 18, 19, 24 Thus, the resolved. Like the clean Si(001) surface, the surface bulk-like Si(001) surface. Close inspection of fig. 3a the clean Si(001) surface show bean-shaped dimers having only a single maximum in the center of the Si=Si dimers and are attributed to P-P dimers atop a chemical identity (and hence, different electronic contrast, under identical conditions STM images of dimers in fig. 3a are clearly distinguishable from unit cell of these surfaces has a (2x1) periodicity, internal structure of the surface unit cell can be states), the STM images show that each (2x1) unit also shows that the white protrusions are always negative sample bias (probing occupied surface cell is clearly resolved into two protrusions. In function of coverage and temperature, some of In fig. 2b and particularly in fig. 3a, the primarily from a delocalized π bonding orbital located one side or the other of the dimer (50% unit cell, due to the fact that electrons tunnel structures). Through controlled studies as a but with an internal structure substantially different from that of the clean surface. At

which are described below, we conclude that the "bright" sites observed in fig. 3 arise from Si-P heterodimers in which the lower (darker) atom is phosphorus and the higher (brighter) atom is silicon. Thus, the phosphorus-terminated surfaces observed in fig. 1-3 consist primarily of P-P dimers, with occasional P-Si dimers.

minimum 0.10 Å deep, and that the "bright" features when we did succeed in getting stable tunneling the are about 0.3 Å higher than the surrounding dimers. 3b, which shows the measured height change along difficult to obtain to achieve stable tunneling, even constant-current STM tunneling across the P-P and at currents as low as 20 picoamperes. Furthermore, a line between the unmarked arrowheads in fig. 3a. The line profile shows that P-P dimers give rise to Si-P dimers are shown more quantitatively in fig. location of the unoccupied surface states. Under electronic structure, we also attempted to obtain positive-bias conditions, we found it extremely surfaces at positive bias, to identify the spatial two protrusions 3.2±0.1 Å apart separated by a atomic-resolution images of the P-terminated Since the STM primarily probes the valence The height changes measured under

surface showed very little corrugation. However, we were able to determine that the protrusions observed at negative bias and attributed to the Si "dangling bonds" of Si-P dimers also appear as protrusions at positive bias, consistent with this overall picture.

Figure 4 depicts the structures of Si=Si, P-P, and P-Si dimers atop the Si(001) surface. In order to facilitate the discussion section, we have also included approximate bond lengths based on literature data for dimerized silicon surface and for inorganic silicon-phosphorus compounds, as discussed below.

In order to help identify the nature of the line defects, we also measured corrugation profiles across the line defects, parallel to the dimer rows.

STM images such as fig. 2b show that the line defects appear as if the two dimers have been pulled in toward the bulk and slightly separated, resulting in the formation of a narrow gap between them. Fig. 5 shows the height profile measured across such a defect, measured between the arrows shown in fig.

2b. The most important result to be emphasized here is that the total width of the defect is only two

dimers wide, with the defect having a symmetry plane which passes midway between two dimers of the surface plane, and passes through the secondlayer atom. This structure is included in a schematic fashion in fig. 5 to better illustrate the symmetry of the defects. As will be discussed later, the basic symmetry of the defect can be used to discriminate between structural models for these defects.

We have also characterized the surfaces shown in fig. 1-3 using Auger electron spectroscopy for elemental analysis and using tunneling spectroscopy to probe the valence electronic structure of the surface. In fig. 6a, the Auger spectrum shows a large silicon peak at 92 eV and a large phosphorus peak at 121 eV, with a peak ratio of approximately 0.57, demonstrating a large amount of phosphorus within the outermost 10 Å of the surface, most of which we attribute to phosphorus forming P2 dimers on the surface.

Fig. 6b shows the results of tunneling spectroscopy measurements made on the surfaces shown in fig 1-2 both before and preparation of the phosphorus overlayer as described for fig. 1. For these measurements the sample-tip distance was

to the lone-pair orbitals on the phosphorus atoms, as approximately mid-gap. These results are consistent P-P dimers, we attribute this -1.1 eV electronic state symmetrically distributed about 0 V sample-tip bias, well resolved protrusions separated by 3.2 Å for the about 1.2 Volts. The sharp increase in current near characteristics are symmetrically distributed about than 1.0 Volt and for positive voltages smaller than plunging toward zero for negative voltage smaller simplicity, we plot the logarithm of the tunneling current vs. sample voltage. On the clean surface, surface, fig. 6b shows that the gap has broadened with previous measurements of the clean Si(001) the data shows a gap approximately 0.85 eV wide, density of states at 1.1 V below the Fermi energy. Based on the fact that the STM images reveal two considerably, with the tunneling current rapid regulated using a sample voltage of -1.5 V and a levels of the clean Si=Si dimer; the gap is almost After formation of the P-terminated -1.1 V indicates a sharp increase in the surface occupied π and the unoccupied π^* antibonding which is the energy difference between the tunneling current of 800 picoamperes. For indicating that the Fermi level is pinned depicted in fig. 4. Again, the tunneling surface. 24

zero sample-tip bias, indicating that although phosphorus substituting on a bulk Si lattice site acts as an n-type dopant, the Fermi energy for these phosphorus-passivated surfaces is still pinned approximately mid-gap.

b) Structure and properties of phosphorus overlayers below monolayer coverage

While at high coverage the surface consists primarily of P-P dimers and some Si-P dimers, this relative amounts of Si-P and Si=Si dimers will depend on the local phosphorus coverage. Figures 1-3 show that when the phosphorus coverage is slightly less than unity, the surface essentially consists of an alloy of P-P and Si-P dimers. In order to investigate this alloying in more detailed, we conducted investigations of the surface structure with a lower coverage of phosphorus. These results show that reducing the phosphorus coverage significantly changes the surface morphology, the types of defects present, and the manner in which islands nucleate and growth on the surface.

Figures 7 and 8 show images at successively higher resolution of a Si(001) surface which was prepared by exposing the Si(001) surface to only 1.5

indicated by "DS". Unlike at higher coverage where displacement, resulting in the formation of strongly substantially smaller, however, as in fig. 7 it is quite coverage are "API" and "AP2" antiphase boundaries in fig. 7a shows that there is still substantial silicon islands produced by Si ejection, while in the earlier LD defects with AP defects also changes the manner dimer strings nucleated adjacent to the line defects, observed in fig. 7a and 7b. The replacement of the easy to distinguish the original substrate from the 1-3) PH₃ at 825 Kelvin. The large-scale STM image in fig. 7 each of these dimer strings has nucleated Langmuirs (a factor of 13 lower exposure than fig. anisotropic islands much like those in the earlier The API and AP2 boundaries can be surface. For example, in the lower left region of figure 7a there are several small dimer strings, difference from earlier images, is the complete mages. The amount of silicon displacement is Instead, the only line defects observed at this in which islands nucleate and growth on the substrate terraces. A second, but less obvious pronounced at higher phosphorus coverage. formed (with 50% probability) when islands fig. 1-3 it was difficult to identify island and absence of the line defects which were so intersect.²³

on an AP2 antiphase boundary, just like clean silicon. Thus, the nucleation and growth kinetics occurring on these surfaces appears to be more similar to that of clean silicon than to the P-terminated surface.

random, we counted the number of Si=Si, Si-P, and P-At higher resolution, figure 8a shows that the count the individual atoms. As in the earlier images, dimers are indicated by "Si=Si" in fig. 8b. In order to dimers, and 244 Si-P dimers. For comparison, figure However, at this lower P coverage some Si=Si dimers topmost region), we counted 311 P-P dimers, 13 Si-Si can be observed, giving rise to bean-shaped dimers surface again takes on a mottled appearance due to the presence of silicon-phosphorus dimers. At the most of the white protrusions (Si atoms) are located just as on the clean surface. Two examples of Si=Si 9 shows the expected distributions functions for a terrace of figure 8b (the entire image except the P dimers in images such as fig. 8b. On the upper highest resolution, as in fig. 8b, it is possible to silicon and phosphorus in the surface layer is investigate whether the spatial distribution of to one side of the dimers, not on the center. determine the surface stoichiometry and to

surface containing 866 phosphorus atoms and 270 silicon atoms, paired randomly on a surface to form a random alloy of P-P, Si-P, and Si=Si dimers (shaded curves) compared with these experimental results (heavy vertical lines). The experimental results show that the distribution is not far from what would be expected based on pure statistics. However, there is a slight deviation which we believe to be real, resulting in a larger number of Si-P dimers and smaller numbers of Si=Si and P-P dimers than would be expected on the basis of pure statistics.

c) Partial overlayers produced by evaporation

To determine whether the observed surface structure is dominated by the growth kinetics, we also conducted experiments in which a completely phosphorus-terminated surface was heated to partially desorb phosphorus. Fig. 10a shows a 8.500 x 8,500 Å region of a surface which was prepared by taking a surface which was completely passivated with P-P dimers, like those shown in fig. 1, and subsequently annealing to 916 Kelvin for 60 seconds. The surface again shows formation of large islands and fingerlike extensions and

invaginations similar to those in previous figures. Auger electron spectroscopy, as in fig. 10b, confirms that the surface phosphorus concentration is lower, giving a P/Si peak ratio of 0.23, approximately 40% of the peak ratio of 0.57 obtained for the samples used in fig. 8. Note that due to the very short path length of electrons in the 70-100 eV range, however, the P/Si peak ratio may not be strictly proportional to concentration.

and AP2 boundaries which behave similarly to those observed in pure silicon epitaxy; for example, in the fig. 11a, which might be related to the desorption of nucleation on an AP2 domain boundary. Due to the with Si=Si dimers predominating. High-resolution the outermost surface layer, as indicated by "V" in medium-coverage regime, we find no evidence for images sometimes show small vacancy clusters in and 11b, show that the surface again consists of a random alloy of silicon and phosphorus, but now Higher-resolution images, such as fig. 11a line defects as strain reliefs, instead finding API prepared in this way, we are unable to obtain phosphorus from the suface. Similarly to the presence of significant disorder in surfaces lower region of fig. 11a we observe island

Si-P, and some P-P dimers on this surface, indicating kinetically-limited process, but most likely describes between Si-P dimers due to charge transfer from the reliable counting statistics of the number of dimers of each type. However, visual inspection of fig. 11b shows the presence of significant numbers of Si=Si, typically alternates direction along a dimer row) or P atoms to its Si dimer partner. Adsorbate-adsorbate Freeman for As and Sb adatoms on Si(001), but for a appear to be some preferential ordering of the P-Si coverages in fig. 1-3 and 7-8 does not result from a dimers along a given dimer row, with a spacing of the equilibrium distribution of Si=Si, Si-P, and P-P longer-range interactions between dimers which 2a0 between P-Si dimer units. This suggests some might arise from direct electrostatic interactions that the surface remains a nearly random alloy dimers. Just above center in fig. 11b there does even when prepared by this method. Thus, we conclude that the alloying observed at higher might be associated with dimer tilting (which interactions have been treated by Tang and signficantly different geometry.

DISCUSSION

a) Geometric and Electronic Properties of P. P and Si-P dimers on Si(001)

found that the PH3 molecule adsorbed directly on top passivation of the surface can readily be understood based on their electronic structure. As a member of higher temperatures where the PH3 dissociates and group V of the periodic table, phosphorus typically room-temperature adsorption of PH3 on Si(001), we coordination number of 3, although compounds in can substitute into the crystal lattice. Our present possible. In a previous paper ²⁶ investigating the giving the P atom a coordination number of 5. At dimer and instead forming two strong P-Si bonds, H desorbs from the surface, the phosphorus atom of a Si=Si dimer, breaking the weak π bond of the phosphorus-phosphorus and phosphorus-silicon which P has a coordination number of 5 are also dimers. The stability of the P-P dimers and their Our results clearly show the formation of results show that this displacement reaction, in forms compounds in which the P atoms has a

which a P atom adsorbed on top of the surface displaces Si from a Si=Si dimer and ejects it onto the surface, occurs at nearly random locations on the surface, since we find both phosphorus and silicon randomly distributed both on the islands and on the terraces. The final result is that the surface consists primarily of P-P and Si-P dimers.

We first consider the structure of the dimers. On the Si(001) surface, the (1x1) unit cell has dimensions of 3.84 Å x 3.84 Å. The normal Si=Si dimers are formally considered to have a double bond, due to the presence of a strong σ bond and a weak π bond; this bond formation results in a (2x1) unit cell in which the two Si atoms within a dimer are separated by 2.30 Å²⁷. Interaction with chemical species such as hydrogen occurs by breaking the weak π bond, leaving the σ bond intact, and forming two new bonds to the adsorbate; because the strong σ bond remains intact, the Si-Si din 2r bond lengthens only slightly, to 2.51 Å.²⁸

Phosphorus typically forms chemical compounds through sp³ hybridization, such that the three bonds are formed at three legs of a tetrahedron, with the fourth leg occupied by the

which is slightly shorter than the 2.30 Å dimer bond studies show that the rotational barrier between the general shape are found, including P2H4 (Rpp=2.11 and P2(CH3)4 (RPP=2.19 Å), 32 all of which show P-P compounds H3Si-PH2 and (H3Si)3P, in which the Sidistance of 2.11 Å ²⁹ and the hydrogen atoms found mirror plane containing the P-P bond). Replacing the four hydrogen atoms of this molecule with four involve a P-P bond distance of approximately 2.2 Å, length of the clean Si(001) surface. Likewise, Si-P in a "gauche" configuration (on the same side of a only P2H4 adopts the gauche configuration and all Å), 29 PF₂PH₂ (RPp=2.22 Å), 30 P₂I4 (Rpp=2.21 Å), 31 molecule. In the compound P2H4, for example, the bond lengths comparable to the value of 2.21 Å observed for P4 vapor.³³ While in the solid phase steric repulsion between the subsitutuent groups, trans- and gauche- configurations is small. 31, 32 number of other phosphorus compounds of this P₂ atoms are bonded to one another with a bond produces a dimerized, P2-terminated surface. A bond lengths can be estimated from data on the Si atoms of the bulk-terminated Si(001) surface others adopt a trans configuration to minimize Thus, we expect that a P-P dimer on Si(001) will "lone pair" electrons, similar to the ammonia

clean Si(001)-(2x1) For comparison, the As-As dimer P bond lengths are 2.249 and 2.248 Å, respectively. ³⁴ Si(001)-(2x1)H monohydride surface. Since both the length and just slightly longer than the length of a than the Si-Si bond lengths in the bulk (2.35 Å) and also shorter than the 2.41 Å bond length between a determined by Chabal and Raghavachari²⁸ for the These Si-P bond lengths are again slightly smaller for silicon dimers, we expect that the phosphorussingle Si-Si dimer bond, measured at 2.51 Å for the P-P dimer bond and the P-Si dimer-substrate bond are shorter than the corresponding bond lengths terminated surface is in tensile stress relative to bond length in As-terminated Si(001) is 2.55 Å, 27 dimerized Si atom and the underlying Si surface considerably longer than the Si=Si dimer bond Si(001)-(2x1)H monohydride structure.

In addition to the role of bond lengths in surface energetics, Meade and Vanderbill³⁵ have emphasized the contributions of bond angle strain. Electrons in lone pair orbitals give rise to larger Coulomic repulsion terms than those in chemical bonds, so that the P atoms in P-P dimers should deviate from tetrahedral coordination in such a way as to decrease the Si-P and angle. However, this

bond angle can only be decreased by a concommitant *increase* in the Si-P bond distance compared with Si-Si. The net result is that compared with the Si-Si dimer, the P-P dimer has strain induced by both the shorter bond distance and the strained bond angles, which reinforce one another and lead to a high energy for P-P dimers.

can be understood based on the fact that the Si-P and The electronic properties of these P-P dimers occupied and lowest unoccupied orbitals - i.e., there spectroscopy results indicate lies at 1.1 eV below the separation between maxima in the STM images is 3.2 strong separation between the appropriate highest dimers, the gap is considerably larger. At negative pair" orbitals on the P atoms, which our tunneling Fermi energy. This is supported by the line profile surfaces, images obtained at positive and negative is a large bandgap. In STM studies of clean silicon bias appear quite different 19, 24 because electron unoccupied π^* surface states, which lie relatively bias electrons most likely tunnel out of the "lone close to the Fermi energy. In the case of the P-P P-P bonds are quite strong and likely involve a tunneling occurs through the occupied π and measurements in fig. 3, which show that the

spatial and electronic contrast at positive voltages to attributed to a As-Si backbond. They also found only would expect the lone pair orbitals to be. At positive tunneling and the resulting images show very little used by Becker, et al., who found a similarly large spectroscopy data in fig. 6 indicates that the current contrast. This interpretation is consistent with that levels; instead, electrons likely tunnel directly into the conduction band and therefore show no atomic increasing voltage compared with that at negative occupied state near -1.6 V attributed to the arsenic bias, we noted earlier that it is difficult to achieve between P atoms and near the location where one contrast for P-P dimers. Likewise, the tunneling the absence of any P-derived unoccupied energy voltage and does not show a sharp "turn-on" at a Å, significantly larger than the 2.2 Å separation "lone-pair" and an unoccupied state near +1.4 V characteristic voltage. We attribute the lack of bandgap for As-As dimers on Si(001), with an at positive bias increases quite slowly with weak contrast at positive sample voltages.

Although at first the formation of a P-Si dimer with an unsaturated "dangling bond" might be considered unfavorable, our results clearly show

show that it is not energetically unfavorable to form universal gas constant (96,487 Joule mole-1 K-1) and our measurements indicate that the P-Si bond is 0.05 a free energy change of approximately 11 kJ/mole, T is absolute temperature, and if we assume that the we arbitrarily assume to be 500 Kelvin, this implies equilibrium constant values, the data above clearly or 0.1 eV per dimer (0.05 eV per P-Si dimer). Thus, equilibrium is "frozen in" at a temperature which eV more stable than the average energy of a Si=Si measurements produces an equilibrium constant disproportionation reaction 2 (P-Si) → Si=Si + P-P, counting statistics are needed to obtain accurate a P-Si dimer, and it might even be energetically relationship ΔG =-RT In (Keq) where R is the dimer and a P-P dimer. While more detailed substituting in the results of our counting Keq of 0.07. If we use the thermodynamic that this is not true. If we consider the

In the case of a P-Si dimer, the phosphorus atom is again able to achieve a favorable bonding configuration, but the silicon atom is left with a total of three bonds and a fourth "dangling bond". Because this dangling bond is only partially

nvestigations by Yu and Oshiyama¹⁵ have predicted the existence of a stable Si-As heterodimer and have the Si-P bond distance and optimize the bond angles tilting of the dimer; dimer tilling could help reduce doping. Their electronic structure calculations for Si-As dimers are also quite consistent with what we they also predicted that at larger negative voltages occupied, electrons can readily tunnel out of these (filled states) and voltages of less than 1 Volt, they the contrast would reverse and the As atom would atoms at positive sample bias. As a result, these Si atoms appear as protrusions both at negative and transfer from the phosphorus lone-pair to the Si Si atoms at negative sample bias and into these Si around both Si and P atoms. 35 Indeed, on Si(001) observe for Si-P dimers- at negative sample bias appear higher, we have not observed a contrast reversal in the Si-P system. We expect that Si-P 'dangling bond" and might also be stabilized by important intermediate in the substitutional As experimental observations. Recent theoretical suggested that such heterodimers may play an heterodimers might be stabilized by electron predicted that the STM images would reveal a positive sample bias, in agreement with our protrusion over the location of the Si atom.

electronic contrast, we are not able to establish with inspection of the images in fig. 3a and 8b shows that a point defect or step edge) causing one dimer to tilt, bisecting the Si=Si dimer bond is broken (such as by ultimately producing a zig-zag chain of alternatelythe P-P dimers adjacent to P-Si dimers do not appear tilting of Si-P dimers is small, or else that surface Psince tilting a Si=Si dimer transfers electron density from the lower atom to the upper atom. Because the However, we note that the case of Si=Si dimers, it is tilting dimers which propagates 10-15 Å along the and experimental²¹ results show that tilting and charge transfer are related, dimer row away from the initial site. 18, 20 Close the tilting of one dimer usually induces a tilting P dimers do not respond in the same way as Si=Si geometric and its adjacent neighbor in the opposite direction, certainty whether Si-P dimers are tilted or not. to be buckled; this indicates that either (static) often observed that when the mirror-plane STM images contain a mixture of dimers of the clean surface. both theoretical 36, 37

The formation of large numbers of silicon "dangling bonds" upon phosphorus adsorption is also consistent with a previous study by Yu, et al.,

results show that it more likely arises from the large 17 who used photoemission spectroscopy to study the intrinsic unoccupied surface states (the π^* antibond to one full monolayer. At near monolayer coverage, bond created for each phosphorus atom, it is only at overwhelms the number of Si-P dimers, that doping the 2p state rapidly increased in binding energy by phosphorus coverage reached a critical value close formation of P-Si dimers. Since at coverages below effect as arising from charge transfer to a band of energy of the bulk silicon 2p state as a function of high coverages, where the number of P-P dimers 0.5 monolayer there is approximately 1 dangling conduction band. They interpreted this pinning phosphorus coverage. They found that the Si 2p unpinned and the near-surface region becomes band) of the clean Si=Si dimers. However, our 0.45 eV, indicating that the surface becomes be achieved in the near-surface region. number of Si dangling bonds created by the state remained constant in energy until the doped n-type with the Fermi level near the

b) Surface Morphology and Defect Structure

As noted above, one distinguishing feature between the images obtained at high P coverage (near 1 monolayer) and those obtained at lower coverage is that at high coverage, the surface contains a large number of line defects which cut perpendicular to the dimer rows. In a previous study of phosphorus on Si(001) using low-energy electron diffraction, Yu and co-workers⁵, 16, 17 found that the (2x1) reconstruction was retained upon formation of a phosphorus overlayer, but the LEED pattern showed strong streaking. Our images suggest that the streaks most likely arise from the line defects and the extensive islanding which occurs within the LEED electron coherence length of approximately 100 Å.

Line defects on Si(001) have been observed under several circumstances. Niehus, et al.³8 showed that transition metal contamination creates line defects similar to those observed in fig. 1-3, giving rise to (2xn) reconstructions where n can vary between approximately 8 and 12 depending on the amount of Ni contamination; the defect formation was attributed to a strain effects arising from sub-surface nickel.³⁸ We rule out such

contamination effects here based on the fact that the line defects observed in fig. 1-2 disappear when the phosphorus is desorbed from the surface by heating to 1400 Kelvin, while transition metal contamination is usually impossible to remove by such treatments.

the dimer rows meet "head-on" but shifted by 3.85 Å of 3.84 Å (=a0) between the islands perpendicular to low temperatures, Bronikowski, et al. showed that Si island nucleation occurs almost exclusively by first reconstruction intersect, with a translational shift expanding to form elongated Si islands, effectively occur in two ways: antiphase boundaries in which boundaries in which the dimer rows meet side-bythan those observed on the clean Si surface. First, defects visible in fig. 1-3 behave quite differently side are denoted "AP1". In silicon homoepitaxy at burying the AP2 boundary.²³ However, the line the dimer row. Such "antiphase" boundaries can we note that on clean silicon, dimer rows which are denoted "AP2" boundaries, while antiphase A second class of line defects can also be dimensional dimer strings, and subsequently covering up AP2 boundaries to form onecreated when islands exhibiting a (2x1)

intersect without any lateral phase difference-- that overlayers do not arise simply from the intersection nucleate adjacent to the boundary but actually avoid observed in figures 1 and 2. Thus, it is clear that the homoepitaxy. Instead, we believe the line defects in when dimer rows intersect head-on but out of phase Also, whereas as for pure silicon epitaxy the dimers by 1 lattice constant, but also when the dimer rows boundaries, for phosphorus overlayers the islands length, inducing strain in the underlying crystal from the fact that the Si=Si dimer bond length is In contrast, fig. 2 shows that for the substantially different from the P-P dimer bond meet "head-on" simply fill in to form a complete is, the line defects observed for the phosphorus phosphorus overlayer the line defects produced fig. 1-3 are a strain relief mechanism resulting line defects observed here at not the antiphase intrinsic to the phosphorus-overlayer surface. crossing the boundary itself, as can be clearly nucleate efficiently directly on top of the AP2 reconstruction, but they instead appear to be overlayer, and the boundary between them of islands which have out-of-phase (2x1) boundaries observed earlier for silicon disappears.

lattice. Line defects consisting of vacancies then act as a strain relief mechanism.

that ordered "2xn" structures on Si(001) arise from believed to account for many of the missing dimer defects observed on clean Si(001) surfaces. germanium deposition on silicon have shown that Tersoff has recently shown that vacancy defects, results in the formation of line defects which cut several systems. For silicon, it has been proposed allowing dimerization of second-layer atoms in a the 4.3% lattice mismatch between Si and Ge also The formation of defects as a strain relief compressive strain induced by Ge adsorption by direction perpendicular to that of the top-layer response to strain fields created by sub-surface perpendicular to the Ge dimer rows and with a spacing varying with the Ge coverage 10, 11, 42 present either to remove intrinsic strain or in mechanism has been proposed previously for originally proposed by Pandey,³⁹ relieve the atoms. This so-called "\pi-bonded defect" is also 38-41 which are impurities. More recently, investigations of contamination by transition metals or other ordered arrays from defects,

Based on simple symmetry arguments, we can state unequivocally that the line defects observed in bonded defect originally proposed by Pandey and them which looks like at first like a dimer vacancy. The line profile in fig. 5 clearly shows that the total atom, while in the π -bonded defect model the center out the π -bonded defect model as the origin of these trenches. This is further supported by calculations the case of phosphorus overlayers cannot be the π figures 2 and 3 and the line scan in fig. 5 show that width of the line defect, including both "pulled-in" atoms. Therefore, symmetry arguments alone rule between dimers along a dimer row. As a result, the the line defects appear like two dimers which are pulled into the surface, with a slight gap between recently studied by Tersoff. ¹² The STM images in defect lowers the energy by adding tensile stress, Tersoff 12 and Pandey 39 have shown that along a of the strain field produced by π -bonded defects. surface dimers and directly over a second-layer compressive stress), the presence of a π -bonded dimer row (which for a Ge overlayer on Si is in of the defect lies midway between second-layer center of this defect lies midway between two dimers, is 2a0, where a0=3.84 Å is the spacing

thereby reducing the total stress. For phosphorus overlayers, however, we noted above that we expect all bond lengths to be shorter than for the Si=Si dimers, which means that formation of a phosphorsterminated surface places the surface in tensile stress. In that case, adding π -bonded defects would increase both the total stress and the total surface energy. If the major reason for forming defects is to act as a strain relief, then on the P-terminated surface we expect that such defects need to add compressive stress in order to stabilize the surface.

The STM images alone cannot unambiguously determine the structure of these defects because there are several possible models having the correct symmetry. Figure 12 schematically shows the symmetry of the line defects in top and side views. The STM images suggest that the dimers which appear to be pulled in toward the surface are indeed present, and the fact that these defects are observed only at the highest P coverages suggests that they are most likely P-P dimers; these dimers are labeled "I" in fig. 12. The line scan in fig. 5 as well as visual inspection of fig. 2b and 3a shows that, the defect must have a total width of 2a0, with the center of the defect midway between two dimers and directly

the overall symmetry evident in the STM images, the the center of the dimer rows bisecting the P-P dimer bonds. There are many ways of creating line defects Fermi level shift at very high P coverages, where Psubsurface substitutional sites. In order to maintain consistent with these observations, but the evidence While we attributed the absence of any measureable above a second-layer atom as indicated by the dotted dimers predominate, and therefore the line defects defects preserves the mirror plane running along P dimers predominate and the line defects appear. doping effect to the presence of large numbers of creation of a vacancy or a substitution must occur induce any dimer tilling, suggesting that the line either substitution of P for Si or else creation of a cither at all the atoms labeled "2" (and depicted in suggests that the most likely possibilities involve vacancy. We note that Yu and Meyerson found intermediate coverages, it is also possible that P black in fig. 12), or else at the atoms labeled "3". that exposure of Si(001) to PH3 only produced a line in fig. 12. Likewise, the line defects do not locations until very high coverage where P-P atoms do not efficiently enter substitutional may arise from the presence of P atoms in dangling bonds on Si-P heterodimers at

More complex defects structures are also possible, and further work will be required to determine the atomic configuration more definitively.

The line defects appear to be affected by at least two variables: the chemical composition of the surface (P/Si ratio), and the local surface morphology. The influence of chemical composition is quite strong, as images near monolayer phosphorus coverage (as in fig. 1-3) show large numbers of line defects, while images at lower concentration (as in fig. 7-8) do not show line defects at all. Thus, the density of line defects appears to be a strong, non-linear function of the surface phosphorus coverage.

As previously pointed out by Tromp^{8, 9, 11} and Tersoff, ¹² in a simple strain-relief model for a binary system (such as Ge on Si, without allowing for Ge-Si heterodimers), the number density of strain-relieving defects should increase smoothly as the ad-layer coverage is increased. In the case of Ge/Si, for example, strain-relieving defects have a spacing of 12a0 at 1 monolayer coverage, decreasing to 8a0 at 2 monolayer.

10, 11

Applying this idea to the case of arsenic adsorption, where As-As dimers

interfacial stress, however, it has been assumed that plane (with silicon adatoms presumably diffusing to the surface structure consists of a two-phase system, under some conditions but displace silicon from the outermost layer to form As-As dimers in the surface intermediate coverages, while if the curve is linear the system will tend to have regions of adsorbed Assuch as As-As dimers and Si-Si dimers, for example, and that heterodimers such as As-Si dimers are not become favorable to have displacive adsorption at In this and As dimers atop the silicon surface coexisting with form an adsorbed layer on top of the Si substrate energy vs. coverage curve is nonlinear, it can step edges), Tromp showed that if the surface most other models of surfactant growth and the substrate containing Si=Si dimers.

In the Si-P case, our experiments leads us to conclude that the formation of heterodimers such as Si-P can play a significant role in the surface energetics. Our experiments indicate that while P-P dimers add significant tensile stress to the surface (therefore causing line defects), Si-P dimers do not. For a random statistical alloy of Si=Si, P-P, and Si-P dimers, we would expect the fraction of the surface

expense due to defects is small (or zero) at small and intermediate P coverages, but rapidly increases in number as the P occupied by P-P dimers to increase as Θ^2 , where Θ is associated with the disproportionation reaction 2(Pstatistical distribution of Si=Si, Si-P, and P-P dimers increases like Θ^2 . In this case, the density of line coverage approaches unity, in agreement with the fractional surface coverage of phosphorus. we assume that the surface stress is primarily preferential formation of P-Si dimers at the might arise at least partially from the strain experiments. We also note that our counting Si) → Si=Si+P-P is slightly positive, leading to of Si=Si and P-P dimers. The (slightly) nonstatistics show that the free energy change the P-P dimers, then the surface stress also associated with formation of P-P dimers. Even for a surface with a given chemical composition, figures 1 and 2 show that the spacing between line defects is variable. However, there is a consistent trend: the line defects are always closer together on extended terraces and are farther apart on narrow islands and "fingers" extending from larger terraces. Rough counting statistics indicate that the chemical composition of the surface is

terrace with larger number of line defects separated that the defects reduce the overall stress tensor both uniform, so the variation in spacing is likely related measurement of the spacing between line defects on composition of these terraces is essentially the same. the dimer bond direction, which is easily relieved in the smaller dimer islands and fingerlike projections The few residual Si-P dimers can be easily identified by about 3a₀-4a₀. We believe that the variable line resulting in a large decrease in the overall surface the islands yields an average spacing of 7a0, while number might arise from stress buildup parallel to parallel to the dimer rows and perpendicular rows, left region of fig. 1 shows many "fingers", with an to the island morphology, rather than its chemical average spacing of about 7-8 a0, extending from a compared with narrow, long islands thus suggests identical on the two terraces. Likewise, the lower due to their narrow width, but on larger terraces on the lower terrace it is 340. Yet, the chemical increase in line defect density on flat terraces produces a higher density of line defects. This protrusions in the image, which is essentially (and counted) based on the density of white composition. Taking fig. 2a as an example, free energy.

¥e thereby changing the total surface energy, the line kinetics by affecting both the island nucleation and growth. Fig. 1-3 show that island nucleation occurs nucleate on antiphase defects and quickly cover up visible at lower left and upper right) and in fig. 2b, the islands become very long and narrow. Because line defects might also be mobile, it is also possible growth occurs at elevated temperatures where the islands. In general, the islands appear to be more the defect, for the phosphorus overlayers just the defects. Irrespective of the exact mechanism, the STM images reveal that further growth occurs in note that unlike epitaxy on Si(001), where island both dimensions but producing very anisotropic opposite is true. The islands nucleate adjacent to islands, once formed, pin the location of the line diriensional dimer strings parallel to the defect. In addition to acting as a strain relief and defects in fig. 1-3 also play a role in the growth that nucleation occurs randomly and that these line defects, but the growing islands tend to be constrained by the line defects; where the line defects are dense, as in the fig. 1b (line defects adjacent to these defects, first forming onc-

growth of silicon or germanium on Si(001). Both for addition to these factors the additional contributions type "A" step edge (which runs parallel to the dimer process, which forms dimer "strings" along the line dimer rows). 43 As a result, the islands tend to grow 'sticking coefficient" for adatoms approaching the type "B" step edge (which cuts perpendicular to the faster parallel to the dimer rows, resulting in long, boundaries (at low coverage); 2) line defects in the from the 1-dimensional shape of the initial nuclei approximately 3:1. In the P/Si system, we have in and for Ge/Si heteroepitaxy, thermodynamic contribution due to the different sticking coefficient for adatom approaching the combination of two effects. First, the effective anisotropic, with an aspect ratio estimated 3 at anisotropic than islands produced by epitaxial combination of four factors: 1) the nucleation and the "bounding" action of the line defects. rows of the islands) is much smaller than the energies of the A and B step edges; 22, 47 the the overall anisotropy can be attributed to a equilibrium island shape is therefore also defects (at high coverage) or at antiphase anisotropic islands arise from a narrow islands. Additionally, there is a Si homoepitaxy^{22, 43}

underlying substrate layer, which appear to constrain the islands: 3) anisotropies in the sticking coefficients for diffusing atoms which react the two inequivalent types of step edges, and 4) thermodynamic asymmetry resulting from the different energies of the two inequivalent step edges. Further work will be required to determine the relative importance of these factors.

Conclusions

Exposure of Si(001) surfaces to PH3 at elevated formation of large numbers of islands on previously temperatures dramatically changes the morphology and chemical properties of the surface through the CVD processes which use PH3 as a dopant precursor. likely accounts for the passivation of Si surfaces in However, the effect of phosphorus is more complex low density of states near the Fermi energy, which neasurements show that these dimers have a very formation of P-P dimers. Tunneling spectroscopy At full reveal drastic changes in morphology, with the han simple passivation, since the surfaces also in turn results in a low chemical reactivity and monolayer coverage, STM images show the formation of P-P dimers and Si-P dimers.

edges. Line defects are visible on the surface at saturation coverage, but the symmetry of the STM images shows that they cannot be the π -bonded defects which were earlier proposed by Tersoff as a strain relief for Ge growth on Si(001); we attribute the difference to the fact that phosphorus overlayers likely place the surface in tensile stress while Ge places the surface in compressive stress. These line defects also influence the growth kinetics and the ultimate growth morphology both by serving as nucleation sites for island growth and by bounding the lateral extent.

At slightly less than monolayer phosphorus coverage, STM images reveal that the line defects disappear and the surface consists of a large numbers of Si-P dimers coexisting with the P-P dimers. Counting statistics indicate that the surface can be described as a nearly-random alloy of P-P, Si=Si, and Si-P dimers but with a small non-statistical preference for formation of Si-P heterodimers. The chemical makeup of the surface appears to strongly influence the growth kinetics; on the surfaces composed predominantly of Si-P dimers the morphology, nucleation, and growth processes are

similar to clean Si, while near monolayer coverage the surface adopts clearly distinct morphology and shows quite different nucleation and growth behavior. Although the formation of heterodimers has not been widely considered previously, our results suggest that such heterodimers might play an important role in strain relief in overlayer systems and are a likely intermediate in dopant incorporation into the silicon bulk during CVD growth.

Acknowledgments

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Figure Captions:

- terminated Si(001) surface, showing spontaneous island formation and the presence of line defects cutting perpendicular to the dimer rows. "1" denotes highest terrace, "5" the lowest. Vsample=-2.0 V, Itunnel=0.2 nA, Dimensions: 1150 Å x 1200 Å.
- 1b) STM image of P-terminated Si(001) surface. Note the dimer string ("DS") nucleated adjacent to a line defect and other islands at upper right bounded on their long sides by line defects. "1" denotes the highest terrace, "4" the lowest. "2S" denotes double-height step. Vsample=-2.0 V, Itunnel=0.2 nA; Dimensions 675 Å x 710 Å.
- appearance of upper and lower terraces, and nucleation of islands adjacent to line defects. Note the "specked" appearance of the surface due to admixture of P and Si atoms on both the islands and the underlying substrate. Vsample=-2.0 V, Itunnel=0.2 nA. Dimensions 340 Å x 365 Å.

- appearance of line defects and the characteristic appearance of line defects and the characteristic appearance of P-P dimers and P-Si dimers. The highest terrace is labeled "I", the main large terrace labeled "2", and the lowest terrace is labeled "3".

 Vsample=-0.2 V, Itunnel= 0.2 nA; Dimensions 173 Å x 170 Å.
- 3a) Enlarged image from fig. 2b, showing the appearance of P-P dimers as individual atoms, and showing that in Si-P dimers the Si atom appears as high protrusion on one side of the dimer, with 50% probability of being found on either side. Vsample=-2.0 V, Itunnel= 0.2 nA. Dimensions 92 Å x 92 Å.
- 3b) Height profile measured between the arrows indicated in fig. 3a, showing that P=P dimers appear as two protrusions with maxima separated by 3.2 Å, and Si "dangling bonds" on Si-P dimers appear about 0.3 Å higher.
- 4) Structural models for Si=Si, P=P, and Si-P dimers, including top and side views. Bond lengths are taken from the literature as described in text..

- 5) Height profile measured across a strain-induced line defect, measured between the two arrows shown in the lower right of fig. 2b, and demonstrating the symmetry of defect with respect to the crystal lattice. Open circles are P atoms of P-P dimers; shaded circles are P atoms in the dimers which appear pulled into the surface in the STM images.
- 6a) Auger electron spectrum of sample prepared as in fig. 1-3, using 3 keV incident electron energy.
- 6b) Tunneling spectroscopy data obtained on clean Si(001) surface (dark line) and on phosphorusterminated surface (thin line), showing the increased bandgap of the P-terminated surface.
- 7a). Large-scale STM image of surface partially covered by phosphorus, prepared as discussed in text. Again note the formation of large anisotropic islands, but the absence of line defects. Vsample=-2.15 V, Itunnel= 0.2 nA. Dimensions 1250 Å x 1180 Å.
- 7b) Higher-resolution STM image of partially P-terminated surface showing formation of highly anisotropic islands and the "speckled" appearance due to the mixture of Si=Si, Si-P, and P-P dimers.

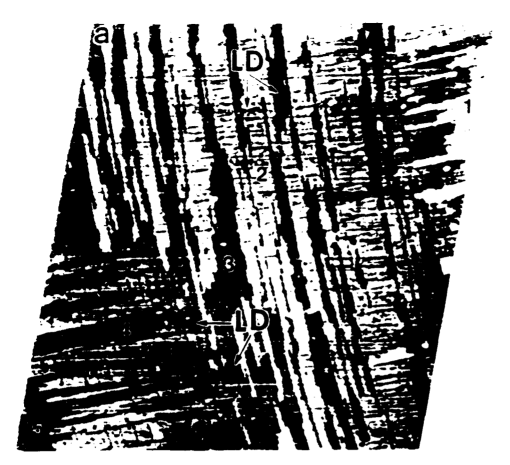
Note that the width of islands are no longer constrained by line defects. Vsample=-2.15 V, Itunne1=0.2 nA. Dimensions 625 Å x 575 Å.

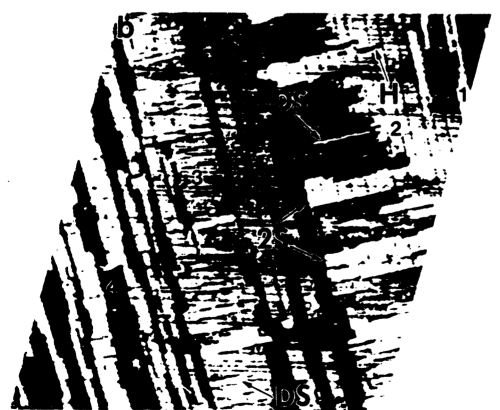
- 8a) High-resolution STM image of partially P-terminated surface showing P-P and Si-P dimers; the chemical composition of the lower and upper terraces is the same. Vsample=-2.2 V, Itunne1=0.2 nA. Dimensions 345 Å x 410 Å.
- 8b) High-resolution image of partially P-terminated surface showing Si=Si, Si-P, and P-P dimers. Some Si=Si dimers are indicated with arrows; Si-P and P-P dimers can be found in numerous locations. There is a step edge near the top of the image. Vsample=-2.2 V, Itunnel=0.2 nA. Dimensions 170 Å x 145 Å.
- omposition obtained by counting the number of Si=Si, Si-P, and P-P dimers on the upper terrace of fig. 8b. Black bars indicate the experimental results, while the shaded graphs show the predicted probability distribution for a completely random alloy containing 866 P atoms and 270 Si atoms.
- 10a) Large-scale STM image of Si(001) surface prepared by partial desorption of phosphorus from

P-terminated surface $V_{sample}=-2.0$ V, $I_{tunnel}=0.1$ nA. Dimensions 8500 Å x 8500 Å.

- 10b) Auger electron spectrum of surface shown in fig. 8, showing that the P concentration is lower but still detectable.
- 11a) STM image of surface prepared by desorption of phosphorus from P-terminated surface showing formation of holes due to phosphorus evaporation.

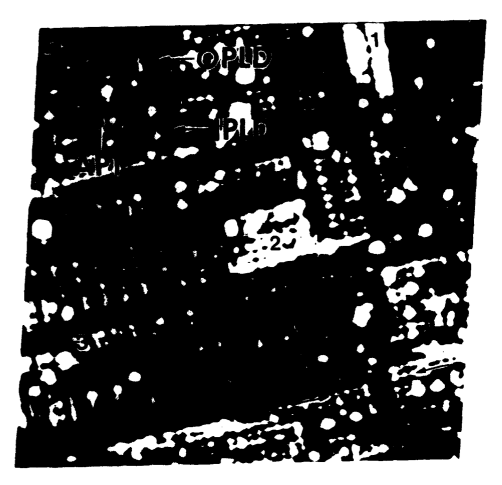
 Vsample=-2.0 V, Itunnel=0.2 nA. Dimensions 341 Å x 261 Å.
- 11b) High-resolution STM image of surface prepared by desorption of phosphorus from P-terminated surface showing P-P dimers, Si-P dimers, and Si-Si dimers. Vsample=-2.0 V, Itunnel=0.2 nA. Dimensions 126 Å x 261 Å.
- 12) Schematic model for line defects. The dotted line indicates the mirror symmetry plane observed in the STM images. Numbered atoms are described in text.



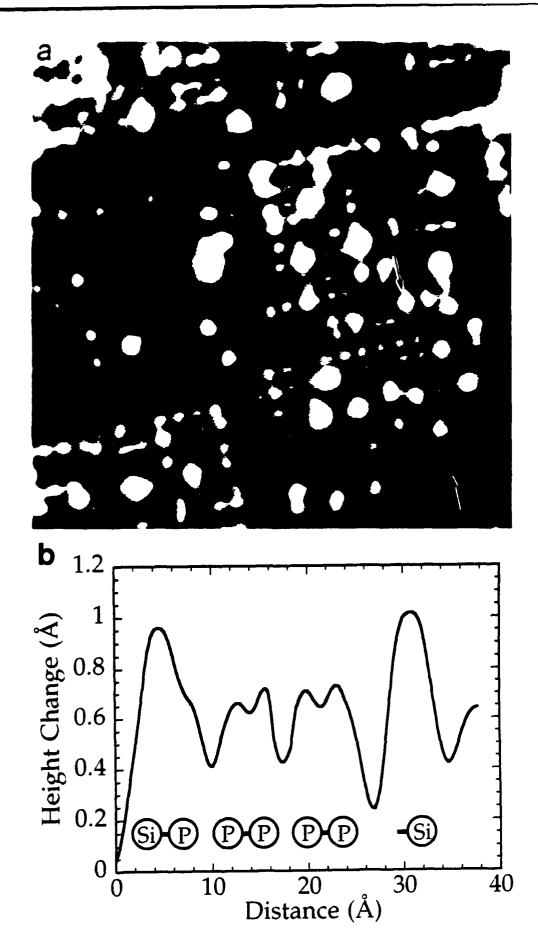


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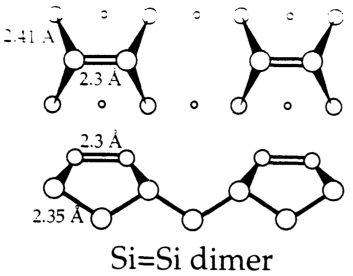


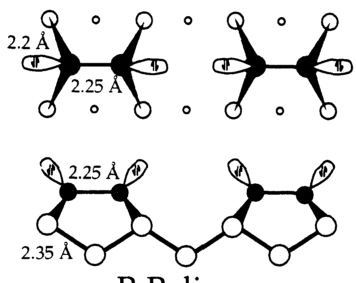


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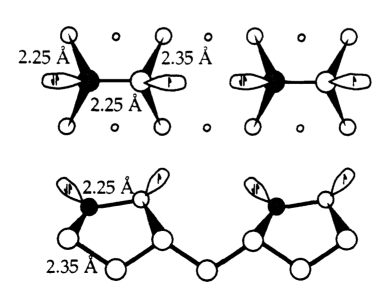


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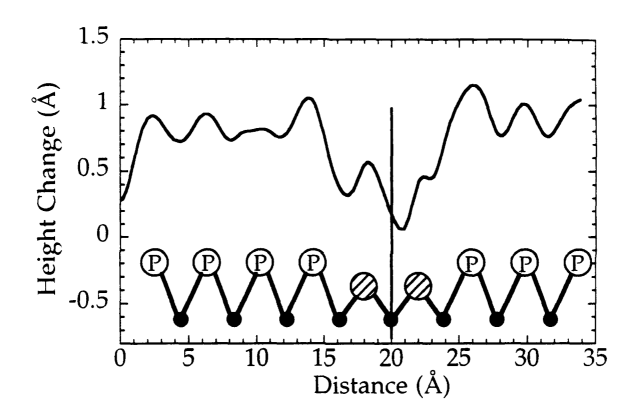


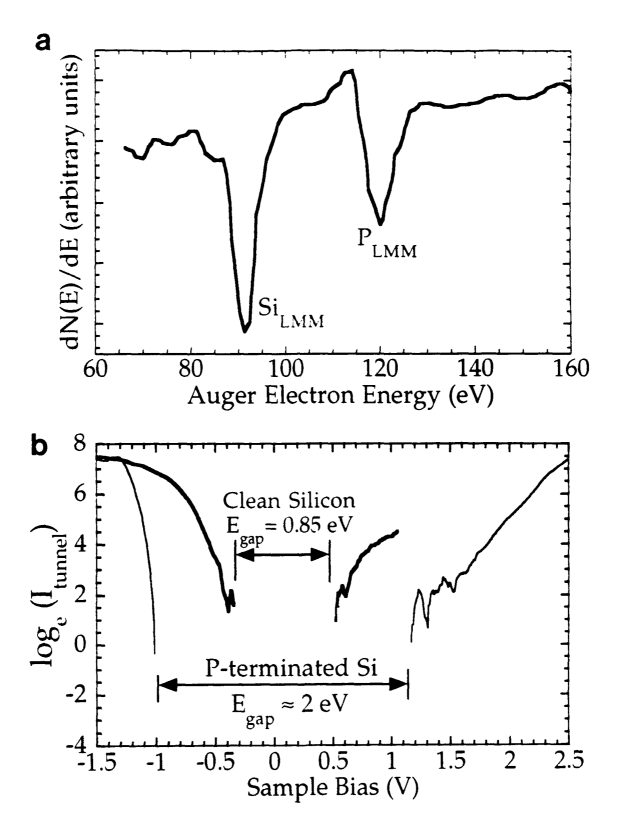


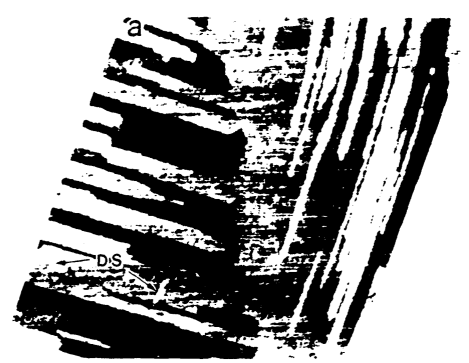
P-P dimer



Si-P dimer

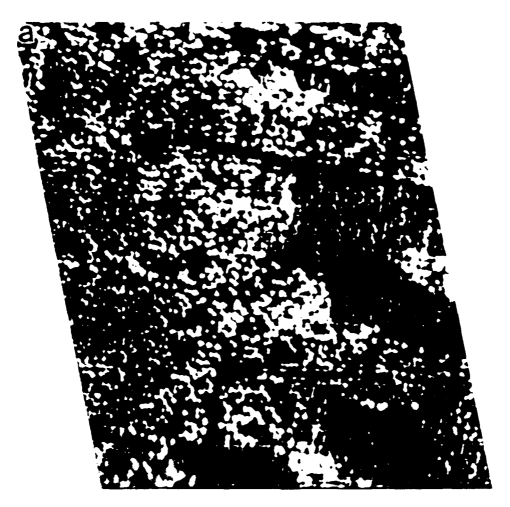


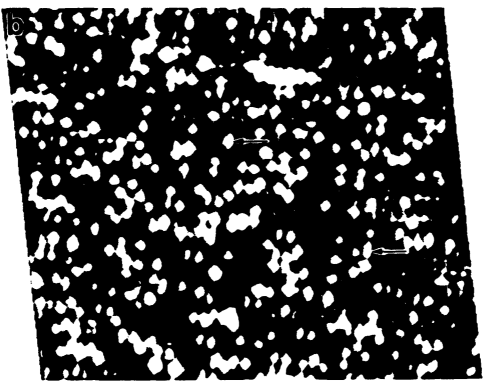


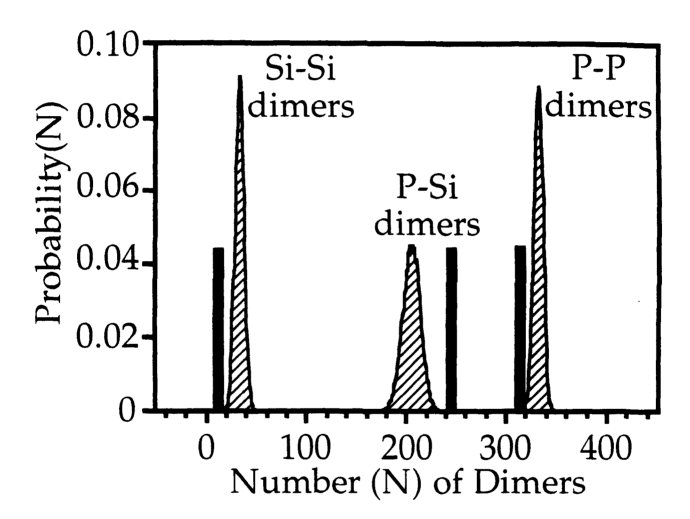




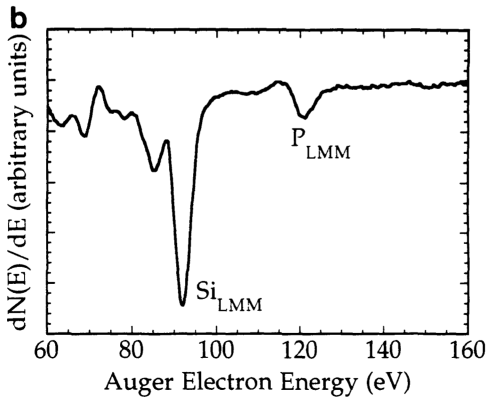
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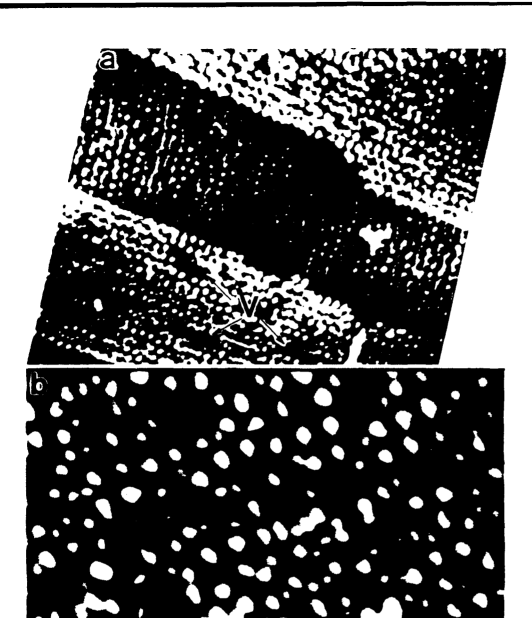




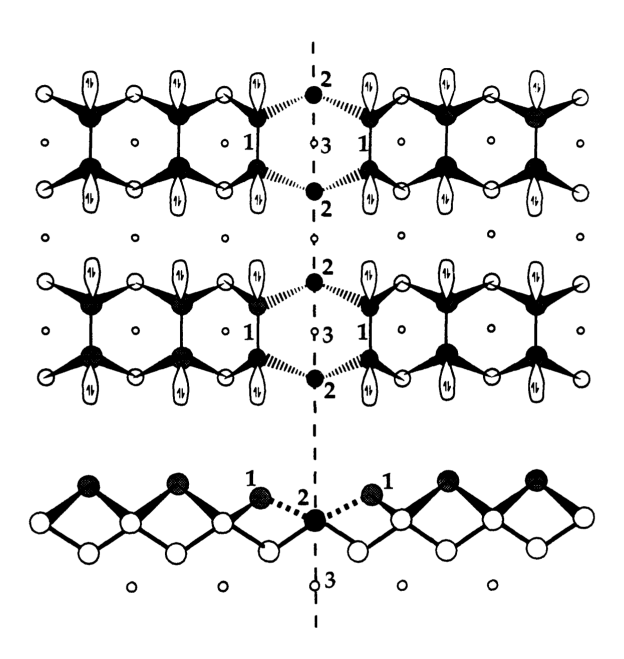








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